HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use Valcyte safely and effectively. See full prescribing information for Valcyte.

Valcyte (valganciclovir hydrochloride) tablets Valcyte (valganciclovir hydrochloride) for oral solution Initial U.S. Approval: 2001

WARNING: HEMATOLOGIC TOXICITY, CARCINOGENICITY, TERATOGENICITY, AND IMPAIRMENT OF FERTILITY

See full prescribing information for complete boxed warning.

- Clinical toxicity of Valcyte, which is metabolized to ganciclovir, includes granulocytopenia, anemia, and thrombocytopenia (5.1)
- In animal studies, ganciclovir was carcinogenic, teratogenic, and caused aspermatogenesis (5.2, 5.3, 5.4)

RECENT MAJOR CHANGES	
Indications and Usage, Pediatric Patients (1.2)	8/2009
Indications and Usage, Limitations of Use (1.3)	8/2009
Dosage and Administration, Pediatric Patients (2.3)	8/2009
Dosage and Administration, Preparation of Valcyte for Oral	
Solution (2.4)	8/2009

- INDICATIONS AND USAGE -

Valcyte is a cytomegalovirus (CMV) nucleoside analogue DNA polymerase inhibitor indicated for:

Adult Patients (1.1)

- Treatment of CMV retinitis in patients with acquired immunodeficiency syndrome (AIDS)
- Prevention of CMV disease in kidney, heart, and kidney-pancreas transplant patients at high risk

Pediatric Patients (1.2)

Prevention of CMV disease in kidney and heart transplant patients at high rick

Limitations of Use (1.3)

- Valcyte is not indicated for use in either adult or pediatric liver transplant patients
- The safety and efficacy of Valcyte have not been established for:
 - Prevention of CMV disease in solid organ transplants other than those indicated
 - Prevention of CMV disease in pediatric solid organ transplant patients < 4 months of age
 - Treatment of congenital CMV disease

DOSAGE AND ADMINISTRATION -

Adult Dosage (2.2)			
Treatment of CMV Retinitis	Induction: 900 mg (two 450 mg tablets) twice a day for 21 days Maintenance: 900 mg (two 450 mg tablets) once a day		
Prevention of CMV Disease	900 mg (two 450 mg tablets) once a day within 10 days of transplantation until 100 days post-transplantation		
Ped	iatric Dosage (2.3)		
Prevention of CMV Disease (4 months to 16 years of age)	Dose once daily according to dosage algorithm (note the calculation of creatinine clearance using a modified Schwartz formula in children 1 to < 2 years of age)		

- Valcyte for oral solution and tablets should be taken with food (2.1).
- Valcyte for oral solution and tablets cannot be substituted for ganciclovir capsules on a one-to-one basis (2.1).
- Adult patients should use Valcyte tablets, not Valcyte for oral solution (2.1).

Adults with renal impairment: Adjust dose based on creatinine clearance.
 For adult patients receiving hemodialysis a dose recommendation cannot be given (2.5, 8.6, 12.3).

DOSAGE FORMS AND STRENGTHS

- Tablets: 450 mg (3)
- Valcyte for Oral Solution: 50 mg/mL (3)

CONTRAINDICATIONS -

Hypersensitivity to valganciclovir or ganciclovir (4)

- WARNINGS AND PRECAUTIONS

- Hematologic effects: Severe leukopenia, neutropenia, anemia, thrombocytopenia, pancytopenia, bone marrow depression, and aplastic anemia have occurred with the use of Valcyte or ganciclovir. Do not administer Valcyte if absolute neutrophil count is < 500 cells/μL, platelet count is < 25,000/μL, or hemoglobin is < 8 g/dL. Use with caution in pre-existing cytopenias and when receiving myelosuppressive drugs or irradiation. Monitor with frequent testing of platelet and complete blood counts (5.1).
- Impairment of fertility: Based on animal studies, Valcyte may cause temporary or permanent inhibition of spermatogenesis (5.2).
- Teratogenesis and mutagenesis: Based on animal studies, Valcyte is
 potentially teratogenic and mutagenic. Women of childbearing potential
 should use contraception during and following treatment and men should
 practice barrier contraception during and following treatment (5.3).
- Acute renal failure: Acute renal failure may occur in elderly patients (with
 or without reduced renal function), patients who receive concomitant
 nephrotoxic drugs, or inadequately hydrated patients. Use with caution in
 elderly patients or those taking nephrotoxic drugs, reduce dosage in patients
 with renal impairment, and monitor renal function (2.5, 8.5, 8.6).

- ADVERSE REACTIONS

- Adult patients: Most common adverse events and laboratory abnormalities (reported in at least one indication by ≥ 20% of patients) are diarrhea, pyrexia, nausea, tremor, neutropenia, anemia, graft rejection, thrombocytopenia, and vomiting (6.1).
- Pediatric patients: Most common adverse events and laboratory abnormalities (reported in > 10% of pediatric solid organ transplant recipients) are diarrhea, pyrexia, hypertension, upper respiratory tract infection, vomiting, anemia, neutropenia, constipation, nausea, and cough (6.2).

To report SUSPECTED ADVERSE REACTIONS, contact Roche at 1-800-526-6367 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

To report SUSPECTED ADVERSE REACTIONS, contact at or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch

- DRUG INTERACTIONS

- Zidovudine: Potential to cause neutropenia and anemia. Monitor with frequent tests of white blood cell counts with differential and hemoglobin levels (7).
- Probenecid: May increase ganciclovir levels. Monitor for evidence of ganciclovir toxicity (7).
- Mycophenolate mofetil (MMF): May increase ganciclovir concentrations and levels of MMF metabolites in patients with renal impairment. Monitor for ganciclovir and MMF toxicity (7).
- Didanosine: May increase didanosine concentrations. Monitor for didanosine toxicity (7).

USE IN SPECIFIC POPULATIONS

- Pregnancy: Based on animal data, Valcyte may cause fetal harm (8.1).
- Nursing mothers: May cause adverse events in nursing infants. Discontinue drug or nursing, taking into consideration the importance of drug to mother (8.3).

See 17 for PATIENT COUNSELING INFORMATION and FDA-approved patient labeling

Revised: 08/2009

TERATOGENICITY, AND IMPAIRMENT OF FERTILITY 1 INDICATIONS AND USAGE

- 1.1 Adult Patients
- 1.2 Pediatric Patients
- 1.3 Limitations of Use

2 DOSAGE AND ADMINISTRATION

- 2.1 General Dosing Information
- 2.2 Adult Patients
- 2.3 Pediatric Patients
- 2.4 Preparation of Valcyte for Oral Solution
- 2.5 Renal Impairment
- 2.6 Handling and Disposal

3 DOSAGE FORMS AND STRENGTHS

4 CONTRAINDICATIONS

5 WARNINGS AND PRECAUTIONS

- 5.1 Hematologic Effects
- 5.2 Impairment of Fertility
- 5.3 Teratogenesis and Mutagenesis
- 5.4 Carcinogenesis
- 5.5 Acute Renal Failure

6 ADVERSE REACTIONS

- 6.1 Clinical Trial Experience in Adult Patients
- 6.2 Clinical Trial Experience in Pediatric Patients
- 6.3 Postmarketing Experience

7 DRUG INTERACTIONS

8 USE IN SPECIFIC POPULATIONS

- 8.1 Pregnancy
- 8.3 Nursing Mothers
- 8.4 Pediatric Use
- 8.5 Geriatric Use
- 8.6 Renal Impairment
- 8.7 Hepatic Impairment

10 OVERDOSAGE

11 DESCRIPTION

12 CLINICAL PHARMACOLOGY

- 12.1 Mechanism of Action
- 12.3 Pharmacokinetics
- 12.4 Virology

13 NONCLINICAL TOXICOLOGY

- 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
- 13.3 Reproductive and Developmental Toxicology

14 CLINICAL STUDIES

- 14.1 Adult Patients
- 14.2 Pediatric Patients

15 REFERENCES

16 HOW SUPPLIED/STORAGE AND HANDLING

17 PATIENT COUNSELING INFORMATION

^{*} Sections or subsections omitted from the full prescribing information are not listed

FULL PRESCRIBING INFORMATION

WARNING: HEMATOLOGIC TOXICITY, CARCINOGENICITY, TERATOGENICITY, AND IMPAIRMENT OF FERTILITY

- Clinical toxicity of Valcyte, which is metabolized to ganciclovir, includes granulocytopenia, anemia, and thrombocytopenia [see Warnings and Precautions (5.1)].
- In animal studies, ganciclovir was carcinogenic, teratogenic, and caused aspermatogenesis [see Warnings and Precautions (5.2, 5.3, 5.4)].

1 INDICATIONS AND USAGE

1.1 Adult Patients

<u>Treatment of Cytomegalovirus (CMV) Retinitis:</u> Valcyte tablets are indicated for the treatment of CMV retinitis in patients with acquired immunodeficiency syndrome (AIDS) [see Clinical Studies (14.1)].

<u>Prevention of CMV Disease:</u> Valcyte tablets are indicated for the prevention of CMV disease in kidney, heart, and kidney-pancreas transplant patients at high risk (Donor CMV seropositive/Recipient CMV seronegative [D+/R-]) [see Clinical Studies (14.1)].

1.2 Pediatric Patients

<u>Prevention of CMV Disease:</u> Valcyte for oral solution and tablets are indicated for the prevention of CMV disease in kidney and heart transplant patients (4 months to 16 years of age) at high risk [see Clinical Studies (14.2)].

1.3 Limitations of Use

Valcyte is not indicated for use in either adult or pediatric liver transplant patients [see Clinical Studies (14.1, 14.2)] The safety and efficacy of Valcyte have not been established for:

- Prevention of CMV disease in solid organ transplants other than those indicated [see Clinical Studies (14.1, 14.2)]
- Prevention of CMV disease in pediatric solid organ transplant patients < 4 months of age [see Clinical Studies (14.2)]
- Treatment of congenital CMV disease [see Use in Specific Populations (8.4)]

2 DOSAGE AND ADMINISTRATION

2.1 General Dosing Information

- Valcyte for oral solution and tablets should be taken with food [see Clinical Pharmacology (12.3)].
- Valcyte for oral solution (50 mg/mL) must be prepared by the pharmacist prior to dispensing to the patient [see Dosage and Administration (2.4)].
- The bioavailability of ganciclovir from Valcyte is significantly higher than from ganciclovir capsules. Therefore, Valcyte tablets cannot be substituted for ganciclovir capsules on a one-to-one basis [see Clinical Pharmacology (12.3)].
- Adult patients should use Valcyte tablets, not Valcyte for oral solution.

2.2 Adult Patients

Treatment of CMV Retinitis in Adult Patients With Normal Renal Function:

- Induction: The recommended dose is 900 mg (two 450 mg tablets) twice a day for 21 days.
- Maintenance: Following induction treatment, or in adult patients with inactive CMV retinitis, the recommended dose is 900 mg (two 450 mg tablets) once a day.

<u>Prevention of CMV Disease:</u> For adult patients who have received a kidney, heart, or kidney-pancreas transplant, the recommended dose is 900 mg (two 450 mg tablets) once a day starting within 10 days of transplantation until 100 days post-transplantation.

2.3 Pediatric Patients

<u>Prevention of CMV Disease:</u> For pediatric patients 4 months to 16 years of age who have received a kidney or heart transplant, the recommended once daily dose of Valcyte starting within 10 days of transplantation until 100 days post-transplantation is based on body surface area (BSA) and creatinine clearance (CrCl) derived from a modified Schwartz formula, and is calculated using the equation below:

Pediatric Dose (mg) = $7 \times BSA \times CrCl$ (calculated using a modified Schwartz formula), where

$$Mosteller\ BSA\left(m^{2}\right) = \sqrt{\frac{Height\left(cm\right)\times Weight\left(kg\right)}{3600}}$$

$$Schwartz\ Creatinine\ Clearance\left(mL/\min\left/1.73m^{2}\right\right) = \frac{k\times Height\left(cm\right)}{Senson\ Creatinine\left(mg/dL\right)}$$

where k =

0.45 for patients aged < 1 year,

0.45 for patients aged 1 to < 2 years (note k value is 0.45 instead of the typical value of 0.55),

0.55 for boys aged 2 to < 13 years and girls aged 2 to 16 years, and

0.7 for boys aged 13 to 16 years.

All calculated doses should be rounded to the nearest 25 mg increment for the actual deliverable dose. If the calculated dose exceeds 900 mg, a maximum dose of 900 mg should be administered. Valcyte for oral solution is the preferred formulation since it provides the ability to administer a dose calculated according to the formula above; however, Valcyte tablets may be used if the calculated doses are within 10% of available tablet strength (450 mg). For example, if the calculated dose is between 405 mg and 495 mg, one 450 mg tablet may be taken.

2.4 Preparation of Valcyte for Oral Solution

Prior to dispensing to the patient, Valcyte for oral solution must be prepared by the pharmacist as follows [see How Supplied/Storage and Handling (16)]:

- Measure 91 mL of purified water in a graduated cylinder.
- Shake the Valcyte bottle to loosen the powder. Remove the child resistant bottle cap and add approximately half the total amount of water for constitution to the bottle and shake the closed bottle well for about 1 minute. Add the remainder of water and shake the closed bottle well for about 1 minute. This prepared solution contains 50 mg of valganciclovir free base per 1 mL.
- Remove the child resistant bottle cap and push the bottle adapter into the neck of the bottle.
- Close bottle with child resistant bottle cap tightly. This will assure the proper seating of the bottle adapter in the bottle and child resistant status of the cap.
- Store constituted oral solution under refrigeration at 2°C to 8°C (36°F to 46°F) for no longer than 49 days. Do not freeze.
- Write the date of expiration of the constituted oral solution on the bottle label.

The patient package insert, which includes the dosing instructions for patients and 2 oral dispensers, should be dispensed to the patient [see Patient Counseling Information (17)].

2.5 Renal Impairment

Dosage recommendations for adult patients with reduced renal function are provided in **Table 1**. For adult patients on hemodialysis (CrCl <10 mL/min), a dose recommendation for Valcyte cannot be given [see Use in Specific Populations (8.5, 8.6), Clinical Pharmacology (12.3)].

Table 1 Dosage Recommendations for Adult Patients With Impaired Renal Function

Valcyte 450 mg Tablets				
CrCl* (mL/min)	Induction Dose	Maintenance/Prevention Dose		
≥ 60	900 mg twice daily	900 mg once daily		
40 – 59	450 mg twice daily	450 mg once daily		
25 – 39	450 mg once daily	450 mg every 2 days		
10 – 24	450 mg every 2 days	450 mg twice weekly		
< 10 (on hemodialysis)	not recommended	not recommended		

*An estimated creatinine clearance is calculated from serum creatinine by the following formulas:

For males =
$$\frac{(140 - age [years]) \times (body weight [kg])}{(72) \times (serum creatinine [mg/dL])}$$

For females = $0.85 \times \text{male value}$

Dosing in pediatric patients with renal impairment can be done using the recommended equations because CrCl is a component in the calculation [see Dosage and Administration (2.3)].

2.6 Handling and Disposal

Caution should be exercised in the handling of Valcyte tablets and Valcyte for oral solution. Tablets should not be broken or crushed. Because valganciclovir is considered a potential teratogen and carcinogen in humans, caution should be observed in handling broken tablets, the powder for oral solution, and the constituted oral solution [see Warnings and Precautions (5.3, 5.4)]. Avoid direct contact with broken or crushed tablets, the powder for oral solution, and the constituted oral solution with skin or mucous membranes. If such contact occurs, wash thoroughly with soap and water, and rinse eyes thoroughly with plain water.

Because ganciclovir shares some of the properties of antitumor agents (i.e., carcinogenicity and mutagenicity), consideration should be given to handling and disposal according to guidelines issued for antineoplastic drugs. Several guidelines on this subject have been published. However, there is no general agreement that all of the procedures recommended in the guidelines are necessary or appropriate [see References (15)].

3 DOSAGE FORMS AND STRENGTHS

Valcyte Tablets

450 mg, pink, convex oval tablets with "VGC" on one side and "450" on the other side.

Valcyte for Oral Solution

50 mg/mL, supplied as a white to slightly yellow powder for constitution, forming a colorless to brownish yellow tutti-frutti flavored solution. Available in glass bottles containing approximately 100 mL of solution after constitution. Valcyte for oral solution must be constituted by the pharmacist prior to dispensing to the patient [see Dosage and Administration (2.4)].

4 CONTRAINDICATIONS

Valcyte is contraindicated in patients who have had a demonstrated clinically significant hypersensitivity reaction (e.g., anaphylaxis) to valganciclovir, ganciclovir, or any component of the formulation [see Adverse Reactions (6.1)].

5 WARNINGS AND PRECAUTIONS

5.1 Hematologic Effects

Severe leukopenia, neutropenia, anemia, thrombocytopenia, pancytopenia, bone marrow aplasia, and aplastic anemia have been reported in patients treated with Valcyte or ganciclovir. Valcyte should not be administered if the absolute neutrophil count is less than 500 cells/μL, the platelet count is less than 25,000/μL, or the hemoglobin is less than 8 g/dL. Valcyte should also be used with caution in patients with pre-existing cytopenias, or who have received or who are receiving myelosuppressive drugs or irradiation. Cytopenia may occur at any time during treatment and may worsen with continued dosing. Cell counts usually begin to recover within 3 to 7 days after discontinuing drug.

Due to the frequency of neutropenia, anemia, and thrombocytopenia in patients receiving Valcyte [see Adverse Reactions (6.1, 6.2)], complete blood counts with differential and platelet counts should be performed frequently, especially in patients in whom ganciclovir or other nucleoside analogues have previously resulted in leukopenia, or in whom neutrophil counts are less than 1000 cells/µL at the beginning of treatment. Increased monitoring for cytopenias may be warranted if therapy with oral ganciclovir is changed to Valcyte, because of increased plasma concentrations of ganciclovir after Valcyte administration [see Clinical Pharmacology (12.3)].

5.2 Impairment of Fertility

Animal data indicate administration of ganciclovir causes inhibition of spermatogenesis and subsequent infertility. These effects were reversible at lower doses but irreversible at higher doses [see Nonclinical Toxicology (13.1)]. In men, Valcyte at the recommended doses may cause temporary or permanent inhibition of spermatogenesis. Animal data also indicate suppression of fertility in females may occur.

5.3 Teratogenesis and Mutagenesis

Animal data indicate ganciclovir is teratogenic and mutagenic. Therefore, Valcyte should be considered to have the potential to cause birth defects and cancers in humans. Women of childbearing potential should be advised to use effective contraception during treatment and for at least 30 days following treatment with Valcyte. Similarly, men should be advised to practice barrier contraception during and for at least 90 days following treatment with Valcyte [see Dosage and Administration (2.6), Use in Specific Populations (8.1), Nonclinical Toxicology (13.1, 13.3)].

5.4 Carcinogenesis

Animal data indicate that administration of ganciclovir is carcinogenic. Valcyte should therefore be considered a potential carcinogen in humans [see Dosage and Administration (2.6), Nonclinical Toxicology (13.1)].

5.5 Acute Renal Failure

Acute renal failure may occur in:

- Elderly patients with or without reduced renal function. Caution should be exercised when administering Valcyte to geriatric patients, and dosage reduction is recommended for those with impaired renal function [see Dosage and Administration (2.5), Use in Specific Populations (8.5, 8.6)].
- Patients receiving potential nephrotoxic drugs. Caution should be exercised when administering Valcyte to patients receiving potential nephrotoxic drugs.
- Patients without adequate hydration. Adequate hydration should be maintained for all patients.

6 ADVERSE REACTIONS

The following serious adverse events are discussed in greater detail in other sections of the labeling:

- Hematologic adverse events [see Boxed Warning, Warnings and Precautions (5.1)]
- Acute renal failure [see Warnings and Precautions (5.5)]

The most common adverse events and laboratory abnormalities reported in at least one indication by $\geq 20\%$ of adult patients treated with Valcyte tablets are diarrhea, pyrexia, nausea, tremor, neutropenia, anemia, graft rejection, thrombocytopenia, and vomiting. The most common reported adverse events and laboratory abnormalities reported in > 10% of pediatric solid organ transplant recipients treated with Valcyte for oral solution or tablets are diarrhea, pyrexia, hypertension, upper respiratory tract infection, vomiting, anemia, neutropenia, constipation, nausea, and cough.

6.1 Clinical Trial Experience in Adult Patients

Valganciclovir, a prodrug of ganciclovir, is rapidly converted to ganciclovir after oral administration. Adverse events known to be associated with ganciclovir usage can therefore be expected to occur with Valcyte.

Because clinical trials are conducted under widely varying conditions, adverse event rates observed in the clinical trials of a drug cannot be directly compared to rates in the clinical trials of another drug and may not reflect rates observed in practice. Treatment of CMV Retinitis in AIDS Patients: In a clinical study for the treatment of CMV retinitis in HIV-infected patients, the adverse events reported by patients receiving Valcyte tablets (n=79) or intravenous ganciclovir (n=79) for 28 days of randomized therapy (21 days induction dose and 7 days maintenance dose), respectively, included diarrhea (16%, 10%), nausea (8%, 14%), headache (9%, 5%), and catheter-related infections (3%, 11%). The incidence of adverse events was similar between the group who received Valcyte tablets and the group who received intravenous ganciclovir, with the exception of catheter-related infections, which occurred with greater frequency in patients randomized to receive intravenous ganciclovir. The frequencies of neutropenia (ANC < 500/µL) were 11% for patients receiving Valcyte tablets compared with 13% for patients receiving intravenous ganciclovir. Anemia (Hgb < 8 g/dL) occurred in 8% of patients in each group. Other laboratory abnormalities occurred with similar frequencies in the two groups.

Adverse events and abnormal laboratory values data are available for 370 patients who received maintenance therapy with Valcyte tablets 900 mg once daily in two open-label clinical trials. Approximately 252 (68%) of these patients received Valcyte tablets for more than nine months (maximum duration was 36 months). **Table 2** and **Table 3** show the pooled adverse event data and abnormal laboratory values from these patients.

Table 2 Pooled Selected Adverse Events Reported in ≥ 5% of Patients who Received Valcyte Tablets Maintenance Therapy for CMV Retinitis

	Patients with CMV Retinitis		
Adverse Events According to Body System	Valcyte Tablets (N=370) %		
Gastrointestinal system			
Diarrhea	41		
Nausea	30		
Vomiting	21		
Abdominal pain	15		
Body as a whole			

Pyrexia	31
Headache	22
Central and peripheral nervous system	
Insomnia	16
Peripheral neuropathy	9
Paresthesia	8
Special senses	
Retinal detachment	15

Table 3 Pooled Laboratory Abnormalities Reported in Patients Who Received Valcyte Tablets Maintenance Therapy for the Treatment of CMV Retinitis

	Patients with CMV Retinitis
Laboratory Abnormalities	Valcyte Tablets (N=370) %
Neutropenia: ANC/μL < 500 500 - < 750 750 - < 1000	19 17 17
Anemia: Hemoglobin g/dL < 6.5 6.5 - < 8.0 8.0 - < 9.5	7 13 16
Thrombocytopenia: Platelets/μL < 25000 25000 - < 50000 50000 - < 100000	4 6 22
Serum Creatinine: mg/dL > 2.5 > 1.5 - 2.5	3 12

Prevention of CMV Disease in Selected Solid Organ Transplantation: Table 4 shows selected adverse events regardless of severity and drug relationship with an incidence of $\geq 5\%$ from a clinical trial (up to 28 days after study treatment) where heart, kidney, kidney-pancreas and liver transplant patients received Valcyte tablets (N=244) or oral ganciclovir (N=126). The majority of the adverse events were of mild or moderate intensity.

Table 4 Percentage of Selected Grades 1-4 Adverse Events Reported in \geq 5% of Patients From a Study of Selected Solid Organ Transplant Patients

Adverse Event	Valcyte Tablets (N=244) %	Oral Ganciclovir (N=126) %	
Diarrhea	30	29	
Tremors	28	25	
Graft rejection	24	30	
Nausea	23	23	
Headache	22	27	
Insomnia	20	16	
Hypertension	18	15	
Vomiting	16	14	
Pyrexia	13	14	

Adverse events not included in **Table 4**, which either occurred at a frequency of $\geq 5\%$ in a clinical study with solid organ transplant patients, or were selected serious adverse events reported in studies with patients with CMV retinitis or in studies with solid organ transplant patients with a frequency of < 5% are listed below.

Allergic reactions: valganciclovir hypersensitivity

Bleeding complications: potentially life-threatening bleeding associated with thrombocytopenia

Central and peripheral nervous system: paresthesia, dizziness (excluding vertigo), convulsion

Gastrointestinal disorders: abdominal pain, constipation, dyspepsia, abdominal distention, ascites

General disorders and administration site disorders: fatigue, pain, edema, peripheral edema, weakness

Hemic system: anemia, neutropenia, thrombocytopenia, pancytopenia, bone marrow depression, aplastic anemia

Hepatobiliary disorders: abnormal hepatic function

Infections and infestations: pharyngitis/nasopharyngitis, upper respiratory tract infection, urinary tract infection, local and systemic infections and sepsis, postoperative wound infection

Injury, poisoning, and procedural complications: postoperative complications, postoperative pain, increased wound drainage, wound dehiscence

Metabolism and nutrition disorders: hyperkalemia, hypokalemia, hypomagnesemia, hyperglycemia, appetite decreased, dehydration, hypophosphatemia, hypocalcemia

Musculoskeletal and connective tissue disorders: back pain, arthralgia, muscle cramps, limb pain

Psychiatric disorders: depression, psychosis, hallucinations, confusion, agitation

Renal and urinary disorders: renal impairment, dysuria, decreased creatinine clearance

Respiratory, thoracic and mediastinal disorders: cough, dyspnea, rhinorrhea, pleural effusion

Skin and subcutaneous tissue disorders: dermatitis, pruritus, acne

Vascular disorders: hypotension

Laboratory abnormalities reported with Valcyte tablets in one study in solid organ transplant patients are listed in **Table 5**.

Table 5 Laboratory Abnormalities Reported in a Study of Selected Solid Organ Transplant Patients*

Laboratory Abnormalities	Valcyte Tablets (N=244) %	Ganciclovir Capsules (N=126) %		
Neutropenia: ANC/μL	5	3		
< 500	3	2		
500 – < 750 750 – < 1000	5	2		
Anemia: Hemoglobin g/dL	1	2		
< 6.5	5	7		
6.5 - < 8.0 8.0 - < 9.5	31	25		
Thrombocytopenia: Platelets/µL				
< 25000	0	2		
25000 - < 50000	1	3		
50000 - < 100000	18	21		
Serum Creatinine: mg/dL	14	21		
> 2.5	45	47		
> 1.5 – 2.5				

^{*}Laboratory abnormalities are those reported by investigators.

6.2 Clinical Trial Experience in Pediatric Patients

Valcyte for oral solution and tablets have been studied in 109 pediatric solid organ transplant patients who were at risk for developing CMV disease (aged 4 months to 16 years) and in 24 neonates with symptomatic congenital CMV disease (aged 8 to 34 days), with

duration of ganciclovir exposure ranging from 2 to 100 days. The overall safety profile was similar in pediatric patients as compared to adult patients. However, the rates of certain adverse events and laboratory abnormalities, such as upper respiratory tract infection, pyrexia, nasopharyngitis, anemia, and neutropenia, were reported more frequently in pediatric patients than in adults [see Use in Specific Populations (8.4), Clinical Studies (14.2)].

6.3 Postmarketing Experience

In general, the adverse events reported during the postmarketing use of Valcyte were similar to those identified during the clinical trials and to those reported during the postmarketing use of ganciclovir. Please also refer to the intravenous ganciclovir product information and ganciclovir capsule product information for more information on postmarketing adverse events associated with ganciclovir.

7 DRUG INTERACTIONS

No in vivo drug-drug interaction studies were conducted with valganciclovir. However, because valganciclovir is rapidly and extensively converted to ganciclovir, drug-drug interactions associated with ganciclovir will be expected for Valcyte. Established and other potentially significant drug interactions conducted with ganciclovir are listed in **Table 6**.

Table 6 Established and Other Potentially Significant Drug Interactions With Ganciclovir

Name of the Concomitant Drug	Change in the Concentration of Ganciclovir or Concomitant Drug	Clinical Comment
Didanosine	↓ Ganciclovir ↑ Didanosine	Patients should be closely monitored for didanosine toxicity
Mycophenolate Mofetil (MMF)	 ⇔ Ganciclovir (in patients with normal renal function) ⇔ MMF (in patients with normal renal function) 	Patients with renal impairment should be monitored carefully as levels of MMF metabolites and ganciclovir may increase
Probenicid	↑ Ganciclovir	Patients taking probenicid and Valcyte should be monitored for evidence of ganciclovir toxicity
Zidovudine	↓ Ganciclovir ↑ Zidovudine	Zidovudine and Valcyte each have the potential to cause neutropenia and anemia.

8 USE IN SPECIFIC POPULATIONS

8.1 Pregnancy

<u>Pregnancy Category C.</u> After oral administration, valganciclovir (prodrug) is converted to ganciclovir (active drug) and, therefore, is expected to have reproductive toxicity effects similar to ganciclovir. There are no adequate and well-controlled studies of valganciclovir or ganciclovir use in pregnant women. In animal studies of ganciclovir, embryo-fetal toxicity and structural malformations occurred. Valganciclovir should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

In animal studies, pregnant mice and rabbits received ganciclovir at doses that produced 2 times the human exposure (based on AUC comparison). Treated rabbits had increased rates of fetal resorption, fetal growth retardation, embryolethality, maternal toxicity, cleft palate, anophthalmia/microphthalmia, aplastic organs (kidney and pancreas), hydrocephaly and brachygnathia. In mice, increased fetal resorptions and embryolethality occurred in the presence of maternal/fetal toxicity.

Daily intravenous doses of approximately 1.7 times the human exposure (based on AUC) administered to female mice prior to mating, during gestation, and during lactation caused hypoplasia of the testes and seminal vesicles in month-old male offspring, as well as pathologic changes in the nonglandular region of the stomach.

Data from an ex-vivo human placental model showed that ganciclovir crosses the human placenta. The transfer occurred by passive diffusion and was not saturable over a concentration range of 1 to 10 mg/mL [see Nonclinical Toxicology (13.3)].

8.3 Nursing Mothers

It is not known whether valganciclovir (prodrug) or ganciclovir (active drug) are excreted in human milk. Because valganciclovir caused granulocytopenia, anemia and thrombocytopenia in clinical trials and ganciclovir was mutagenic and carcinogenic in animal studies, serious adverse events may occur from ganciclovir exposure in nursing infants [see Boxed Warning, Warnings and Precautions (5.4)]. Because of the potential for serious adverse events in nursing infants, a decision should be made whether to discontinue nursing or discontinue drug, taking into consideration the importance of the drug to the mother. The Centers for Disease Control and Prevention recommend that HIV-infected mothers not breast-feed their infants to avoid risking postnatal transmission of HIV.

8.4 Pediatric Use

Valcyte for oral solution and tablets are indicated for the prevention of CMV disease in kidney and heart transplant pediatric patients 4 months to 16 years of age at risk for developing CMV disease [see Indications and Usage (1.2), Dosage and Administration (2.3)]. The use of Valcyte for oral solution and tablets for the prevention of CMV disease in pediatric patients 4 months to 16 years of age with kidney or heart transplant is based on pharmacokinetic, safety, and efficacy data from an open-label trial with oral Valcyte (Valcyte for oral solution or tablets) in pediatric solid organ transplant recipients at risk for developing CMV disease. The results of this study were supported by previous demonstration of efficacy in adult patients [see Adverse Reactions (6.2), Clinical Pharmacology (12.3), Clinical Studies (14.2)].

The safety and efficacy of Valcyte for oral solution and tablets have not been established in children for:

- Prevention of CMV disease in liver transplant patients
- Prevention of CMV disease in solid organ transplants other than those indicated
- Prevention of CMV disease in pediatric solid organ transplant patients < 4 months of age
- Treatment of congenital CMV disease

The pharmacokinetic profile and safety of Valcyte for oral solution in children were studied in two open-label studies. Study 1 was an open-label trial with oral Valcyte (Valcyte for oral solution or tablets) in pediatric solid organ transplant recipients at risk for developing CMV disease [see Clinical Pharmacology (12.3), Clinical Studies (14.2)].

Study 2 was a pharmacokinetic and pharmacodynamic evaluation of Valcyte for oral solution in neonates with congenital CMV infection involving the central nervous system. Twenty-four neonates were enrolled in this study. All patients were treated for 6 weeks with a combination of intravenous ganciclovir 6 mg/kg twice daily and Valcyte for oral solution at doses ranging from 14 mg/kg to 20 mg/kg twice daily. The pharmacokinetic results showed that in infants > 7 days to 3 months of age, a dose of 16 mg/kg twice daily of Valcyte for oral solution provided ganciclovir systemic exposures (median $AUC_{0-12h} = 23.6$ [range 16.8 - 35.5] $\mu g \cdot h/mL$; n = 6) comparable to those obtained in infants up to 3 months from a 6 mg/kg dose of intravenous ganciclovir twice daily ($AUC_{0-12h} = 25.3$ [range 2.4 - 89.7] $\mu g \cdot h/mL$; n = 18) or to the ganciclovir systemic exposures obtained in adults from a 900 mg dose of Valcyte tablets twice daily.

The safety and efficacy of intravenous ganciclovir have not been established for the treatment of congenital CMV infection in infants and no similar disease occurs in adults; therefore, efficacy cannot be extrapolated from intravenous ganciclovir use in adults.

8.5 Geriatric Use

No studies of Valcyte for oral solution or tablets have been conducted in adults older than 65 years of age. Clinical studies of Valcyte did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. In general, dose selection for an elderly patient should be cautious, usually starting at the low end of the dosing range, reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy. Valcyte is known to be substantially excreted by the kidneys, and the risk of toxic reactions to this drug may be greater in patients with impaired renal function. Because elderly patients are more likely to have decreased renal function, care should be taken in dose selection. In addition, renal function should be monitored and dosage adjustments should be made accordingly [see Dosage and Administration (2.5), Warnings and Precautions (5.5), Clinical Pharmacology (12.3)].

8.6 Renal Impairment

Dose reduction is recommended when administering Valcyte to patients with renal impairment [see Dosage and Administration (2.5), Warnings and Precautions (5.5)].

For adult patients on hemodialysis (CrCl <10 mL/min) Valcyte tablets should not be used. Adult hemodialysis patients should use ganciclovir in accordance with the dose-reduction algorithm cited in the Cytovene®-IV and ganciclovir capsules complete product information section on DOSAGE AND ADMINISTRATION: Renal Impairment [see Dosage and Administration (2.5) and Clinical Pharmacology (12.3)].

8.7 Hepatic Impairment

The safety and efficacy of Valcyte have not been studied in patients with hepatic impairment.

10 OVERDOSAGE

<u>Experience With Valcyte Tablets:</u> One adult developed fatal bone marrow depression (medullary aplasia) after several days of dosing that was at least 10-fold greater than recommended for the patient's estimated degree of renal impairment.

An overdose of Valcyte could also possibly result in increased renal toxicity [see Dosage and Administration (2.5), Use in Specific Populations (8.6)].

Because ganciclovir is dialyzable, dialysis may be useful in reducing serum concentrations in patients who have received an overdose of Valcyte [see Clinical Pharmacology (12.3)]. Adequate hydration should be maintained. The use of hematopoietic growth factors should be considered [see Clinical Pharmacology (12.3)].

<u>Experience With Intravenous Ganciclovir:</u> Reports of overdoses with intravenous ganciclovir have been received from clinical trials and during postmarketing experience. The majority of patients experienced one or more of the following adverse events:

Hematological toxicity: pancytopenia, bone marrow depression, medullary aplasia, leukopenia, neutropenia, granulocytopenia *Hepatotoxicity:* hepatitis, liver function disorder

Renal toxicity: worsening of hematuria in a patient with pre-existing renal impairment, acute renal failure, elevated creatinine *Gastrointestinal toxicity:* abdominal pain, diarrhea, vomiting

Neurotoxicity: generalized tremor, convulsion

11 DESCRIPTION

Valcyte contains valganciclovir hydrochloride (valganciclovir HCl), a hydrochloride salt of the L-valyl ester of ganciclovir that exists as a mixture of two diastereomers. Ganciclovir is a synthetic guanine derivative active against cytomegalovirus (CMV).

Valcyte is available as a 450 mg tablet for oral administration. Each tablet contains 496.3 mg of valganciclovir HCl (corresponding to 450 mg of valganciclovir), and the inactive ingredients microcrystalline cellulose, povidone K-30, crospovidone and stearic acid. The film-coat applied to the tablets contains Opadry Pink[®].

Valcyte is also available as a powder for oral solution, which when constituted with water as directed contains 50 mg/mL valganciclovir free base. The inactive ingredients of Valcyte for oral solution are sodium benzoate, fumaric acid, povidone K-30, sodium saccharin, mannitol and tutti-frutti flavoring.

Valganciclovir HCl is a white to off-white crystalline powder with a molecular formula of C14H22N6O5•HCl and a molecular weight of 390.83. The chemical name for valganciclovir HCl is L-Valine, 2-[(2-amino-1,6-dihydro-6-oxo-9H-purin-9-yl)methoxy]-3-hydroxypropyl ester, monohydrochloride. Valganciclovir HCl is a polar hydrophilic compound with a solubility of 70 mg/mL in water at 25°C at a pH of 7.0 and an n-octanol/water partition coefficient of 0.0095 at pH 7.0. The pKa for valganciclovir HCl is 7.6. The chemical structure of valganciclovir HCl is:

All doses in this insert are specified in terms of valganciclovir.

12 CLINICAL PHARMACOLOGY

12.1 Mechanism of Action

Valganciclovir is an antiviral drug [see Clinical Pharmacology (12.4)].

12.3 Pharmacokinetics

Because the major elimination pathway for ganciclovir is renal, dosage reductions according to creatinine clearance are required for Valcyte tablets and Valcyte for oral solution [see Dosage and Administration (2.5)].

<u>Pharmacokinetics in adults:</u> The pharmacokinetics of valganciclovir and ganciclovir after administration of valganciclovir tablets have been evaluated in HIV- and CMV-seropositive patients, patients with AIDS and CMV retinitis, and in solid organ transplant patients.

The ganciclovir pharmacokinetic parameters following administration of 900 mg Valcyte tablets and 5 mg/kg intravenous ganciclovir and 1000 mg three times daily oral ganciclovir in HIV-positive/CMV-positive patients are summarized in **Table 7**.

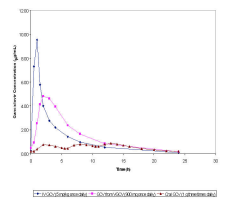
Table 7 Mean Ganciclovir Pharmacokinetic* Measures in Healthy Volunteers and HIV-positive/CMV-positive Adults at Maintenance Dosage

Formulation	Valcyte Tablets	Intravenous Ganciclovir	Ganciclovir Capsules		
Dosage	900 mg once daily with food	5 mg/kg once daily	1000 mg three times daily with food		
AUC _{0-24h} (μg·h/mL)	29.1 ± 9.7 (3 studies, n=57)	26.5 ± 5.9 (4 studies, n=68)	Range of means 12.3 to 19.2 (6 studies, n=94)		
C _{max} (μg/mL)	5.61 ± 1.52 (3 studies, n=58)	9.46 ± 2.02 (4 studies, n=68)	Range of means 0.955 to 1.40 (6 studies, n=94)		
Absolute oral bioavailability (%)	59.4 ± 6.1 (2 studies, n=32)	Not Applicable	Range of means 6.22 ± 1.29 to 8.53 ± 1.53 (2 studies, n=32)		
Elimination half-life (hr)	4.08 ± 0.76 (4 studies, n=73)	3.81 ± 0.71 (4 studies, n=69)	Range of means 3.86 to 5.03 (4 studies, n=61)		
Renal clearance (mL/min/kg)	3.21 ± 0.75 (1 study, n=20)	2.99 ± 0.67 (1 study, n=16)	Range of means 2.67 to 3.98 (3 studies, n=30)		

*Data were obtained from single and multiple dose studies in healthy volunteers, HIV-positive patients, and HIV-positive/CMV-positive patients with and without retinitis. Patients with CMV retinitis tended to have higher ganciclovir plasma concentrations than patients without CMV retinitis.

The area under the plasma concentration-time curve (AUC) of ganciclovir administered as Valcyte tablets (900 mg once daily) is comparable to the AUC of ganciclovir after administration of intravenous ganciclovir (5 mg/kg once daily). The C_{max} of ganciclovir following Valcyte administration is 40% lower than the C_{max} following intravenous ganciclovir administration. During maintenance dosing, ganciclovir AUC_{0-24h} and C_{max} following oral ganciclovir administration (1000 mg three times daily) are lower relative to Valcyte and intravenous ganciclovir. The ganciclovir C_{min} following intravenous ganciclovir and Valcyte administration are less than the ganciclovir C_{min} following oral ganciclovir administration. The clinical significance of the differences in ganciclovir pharmacokinetics after administration of Valcyte tablets, ganciclovir capsules, and intravenous ganciclovir is unknown.

Figure 1 Ganciclovir Plasma Concentration Time Profiles in HIV-positive/CMV-positive Patients*



*Plasma concentration-time profiles for ganciclovir (GCV) from valganciclovir (VGCV) and intravenous ganciclovir were obtained from a multiple dose study (n=21 and n=18, respectively) in HIV-positive/CMV-positive patients with CMV retinitis. The plasma concentration-time profile for oral ganciclovir was obtained from a multiple dose study (n=24) in HIV-positive/CMV-positive patients without CMV retinitis.

In solid organ transplant recipients, the mean systemic exposure to ganciclovir was $1.7 \times$ higher following administration of 900 mg Valcyte tablets once daily versus 1000 mg ganciclovir capsules three times daily, when both drugs were administered according to their renal function dosing algorithms. The systemic ganciclovir exposures attained were comparable across kidney, heart and liver transplant recipients based on a population pharmacokinetics evaluation (see **Table 8**).

Table 8 Mean Ganciclovir Pharmacokinetic Measures by Organ Transplant Type

Parameter	Ganciclovir Capsules	Valcyte Tablets 900 mg once daily with food	
Dosage	1000 mg three times daily with food		
Heart Transplant Recipients	N=13	N=17	
AUC_{0-24h} (µg·h/mL)	26.6 ± 11.6	40.2 ± 11.8	
$C_{max} (\mu g/mL)$	1.4 ± 0.5	4.9 ± 1.1	
Elimination half-life (hr)	8.47 ± 2.84	6.58 ± 1.50	
Liver Transplant Recipients	N=33	N=75	
$AUC_{0-24h} (\mu g \cdot h/mL)$	24.9 ± 10.2	46.0 ± 16.1	
$C_{max} (\mu g/mL)$	1.3 ± 0.4	5.4 ± 1.5	
Elimination half-life (hr)	7.68 ± 2.74	6.18 ± 1.42	
Kidney Transplant Recipients [*]	N=36	N=68	
AUC_{0-24h} (µg·h/mL)	31.3 ± 10.3	48.2 ± 14.6	
$C_{max} (\mu g/mL)$	1.5 ± 0.5	5.3 ± 1.5	
Elimination half-life (hr)	9.44 ± 4.37	6.77 ± 1.25	

In a pharmacokinetic study in liver transplant patients, the ganciclovir AUC_{0-24h} achieved with 900 mg valganciclovir was $41.7 \pm 9.9 \, \mu g \cdot h/mL$ (n=28) and the AUC_{0-24h} achieved with the approved dosage of 5 mg/kg intravenous ganciclovir was $48.2 \pm 17.3 \, \mu g \cdot h/mL$ (n=27).

Absorption: Valganciclovir, a prodrug of ganciclovir, is well absorbed from the gastrointestinal tract and rapidly metabolized in the intestinal wall and liver to ganciclovir. The absolute bioavailability of ganciclovir from Valcyte tablets following administration with food was approximately 60% (3 studies, n=18; n=16; n=28). Ganciclovir median T_{max} following administration of 450 mg to 2625 mg Valcyte tablets ranged from 1 to 3 hours. Dose proportionality with respect to ganciclovir AUC following administration of Valcyte tablets was demonstrated only under fed conditions. Systemic exposure to the prodrug, valganciclovir, is transient and low, and the AUC₂₄ and C_{max} values are approximately 1% and 3% of those of ganciclovir, respectively.

Food Effects: When Valcyte tablets were administered with a high fat meal containing approximately 600 total calories (31.1 g fat, 51.6 g carbohydrates and 22.2 g protein) at a dose of 875 mg once daily to 16 HIV-positive subjects, the steady-state ganciclovir AUC increased by 30% (95% CI 12% to 51%), and the C_{max} increased by 14% (95% CI -5% to 36%), without any prolongation in time to peak plasma concentrations (T_{max}). Valcyte should be administered with food [see Dosage and Administration (2.1)].

Distribution: Due to the rapid conversion of valganciclovir to ganciclovir, plasma protein binding of valganciclovir was not determined. Plasma protein binding of ganciclovir is 1% to 2% over concentrations of 0.5 and 51 μ g/mL. When ganciclovir was administered intravenously, the steady-state volume of distribution of ganciclovir was 0.703 \pm 0.134 L/kg (n=69).

After administration of Valcyte tablets, no correlation was observed between ganciclovir AUC and reciprocal weight; oral dosing of Valcyte tablets according to weight is not required.

Metabolism: Valganciclovir is rapidly hydrolyzed to ganciclovir; no other metabolites have been detected. No metabolite of orally administered radiolabeled ganciclovir (1000 mg single dose) accounted for more than 1% to 2% of the radioactivity recovered in the feces or urine.

Elimination: The major route of elimination of valganciclovir is by renal excretion as ganciclovir through glomerular filtration and active tubular secretion. Systemic clearance of intravenously administered ganciclovir was 3.07 ± 0.64 mL/min/kg (n=68) while renal clearance was 2.99 ± 0.67 mL/min/kg (n=16).

The terminal half-life ($t_{1/2}$) of ganciclovir following oral administration of Valcyte tablets to either healthy or HIV-positive/CMV-positive subjects was 4.08 ± 0.76 hours (n=73), and that following administration of intravenous ganciclovir was 3.81 ± 0.71 hours (n=69). In heart, kidney, kidney-pancreas, and liver transplant patients, the terminal elimination half-life of ganciclovir following oral administration of Valcyte was 6.48 ± 1.38 hours, and following oral administration of ganciclovir capsules was 8.56 ± 3.62 .

Specific Populations:

Renal Impairment: The pharmacokinetics of ganciclovir from a single oral dose of 900 mg Valcyte tablets were evaluated in 24 otherwise healthy individuals with renal impairment.

Table 9 Pharma	cokinetics o	f Ganciclov	ir From a	Single Oral Do	se of 900 mg	Valcyte Tablets
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Estimated Creatinine Clearance (mL/min)	N	Apparent Clearance (mL/min) Mean ± SD	AUC _{last} (μg·h/mL) Mean ± SD	Half-life (hours) Mean ± SD
51-70	6	249 ± 99	49.5 ± 22.4	4.85 ± 1.4
21-50	6	136 ± 64	91.9 ± 43.9	10.2 ± 4.4
11-20	6	45 ± 11	223 ± 46	21.8 ± 5.2
≤10	6	12.8 ± 8	366 ± 66	67.5 ± 34

Decreased renal function results in decreased clearance of ganciclovir from valganciclovir, and a corresponding increase in terminal half-life. Therefore, dosage adjustment is required for patients with impaired renal function.

Hemodialysis reduces plasma concentrations of ganciclovir by about 50% following Valcyte administration. Adult patients receiving hemodialysis (CrCl <10 mL/min) cannot use Valcyte tablets because the daily dose of Valcyte tablets required for these patients is less than 450 mg [see Dosage and Administration (2.5) and Use in Specific Populations (8.6)].

Pharmacokinetics in Pediatric Patients: The pharmacokinetics of ganciclovir were evaluated following the administration of valganciclovir in 63 pediatric solid organ transplant patients aged 4 months to 16 years. In this study, patients received oral doses of

valganciclovir (either Valcyte for oral solution or tablets) to produce exposure equivalent to an adult 900 mg dose [see Dosage and Administration (2.3), Adverse Reactions (6.2), Use in Specific Populations (8.4), Clinical Studies (14.2)].

The pharmacokinetics of ganciclovir were similar across organ types and age ranges. Population pharmacokinetic modeling suggested that bioavailability was approximately 60%. Clearance was positively influenced by both body surface area and renal function. The mean total clearance was 5.3 L/hr (88.3 mL/min) for a patient with creatinine clearance of 70.4 mL/min. The mean C_{max} and AUC by age and organ type are listed in **Table 10**.

Table 10 Mean (SD) Pharmacokinetics of Ganciclovir by Age in Pediatric Solid Organ Transplant Patients

	PK Parameter		Age Group in Years	
		≤ 2 (n=2)	> 2 to < 12 (n=10)*,†	≥ 12 (n=19)
Kidney	$AUC_{0-24h} (\mu g \cdot h/mL)$	67.6 (13.0)	55.9 (12.1)	47.8 (12.4)
(N=31)	$C_{max} (\mu g/mL)$	10.4 (0.4)	8.7 (2.1)	7.7 (2.1)
	$t_{1/2}$ (h)	4.5 (1.5)	4.8 (1.0)	6.0 (1.3)
		≤ 2 (n=9)	> 2 to < 12 (n=6)	≥ 12 (n=2)
Liver	$AUC_{0-24h} (\mu g \cdot h/mL)$	69.9 (37.0)	59.4 (8.1)	35.4 (2.8)
(N=17)	$C_{max} (\mu g/mL)$	11.9 (3.7)	9.5 (2.3)	5.5 (1.1)
	$t_{1/2}$ (h)	2.8 (1.5)	3.8 (0.7)	4.4 (0.2)
		≤ 2 (n =6)	> 2 to < 12 (n=2)	≥ 12 (n=4)
Heart	$AUC_{0-24h} (\mu g \cdot h/mL)$	55.4 (22.8)	59.6 (21.0)	60.6 (25.0)
(N=12)	C_{max} (µg/mL)	8.2 (2.5)	12.5 (1.2)	9.5 (3.3)
	t _{1/2} (h)	3.8 (1.7)	2.8 (0.9)	4.9 (0.8)

^{*}There was one subject in this age group who received both a kidney and liver transplant. The pharmacokinetic profile for this subject has not been included in this table as it is not possible to determine whether the effects observed are from the kidney/liver transplant or neither.

Pharmacokinetics in Geriatric Patients: The pharmacokinetic characteristics of Valcyte in elderly patients have not been established. Because elderly individuals frequently have a reduced glomerular filtration rate, renal function should be assessed before and during administration of Valcyte [see Dosage and Administration (2.5)].

Drug Interactions: No in vivo drug-drug interaction studies were conducted with valganciclovir. However, because valganciclovir is rapidly and extensively converted to ganciclovir, interactions associated with ganciclovir will be expected for Valcyte [see Drug Interactions (7)].

Drug-drug interaction studies were conducted in patients with normal renal function. Patients with impaired renal function may have increased concentrations of ganciclovir and the coadministered drug following concomitant administration of Valcyte and drugs excreted by the same pathway as ganciclovir. Therefore, these patients should be closely monitored for toxicity of ganciclovir and the coadministered drug.

Table 11 and **Table 12** provide a listing of established drug interaction studies with ganciclovir. **Table 11** provides the effects of coadministered drug on ganciclovir plasma pharmacokinetic parameters, whereas **Table 12** provides the effects of ganciclovir on plasma pharmacokinetic parameters of co-administered drug.

Table 11 Results of Drug Interaction Studies With Ganciclovir: Effects of Coadministered Drug on Ganciclovir Pharmacokinetic Parameters

Coadministered Drug	Ganciclovir Dosage	N	Ganciclovir Pharmacokinetic (PK) Parameter
Zidovudine 100 mg every 4 hours	1000 mg every 8 hours	12	AUC ↓ 17 ± 25% (range: -52% to 23%)
Probenecid 500 mg every 6 hours	1000 mg every 8 hours		AUC \uparrow 53 \pm 91% (range: -14% to 299%) Ganciclovir renal clearance \downarrow 22 \pm 20% (range: -54% to -4%)

[†]The pharmacokinetic profiles for two subjects in this age group who received kidney transplants have not been included in this table as the data were determined to be non-evaluable.

Mycophenolate Mofetil (MMF) 1.5 g single dose	IV ganciclovir 5 mg/kg single dose	12	No effect on ganciclovir PK parameters observed (patients with normal renal function)
Didanosine 200 mg every 12 hours administered 2 hours before ganciclovir	1000 mg every 8 hours	12	AUC ↓ 21 ± 17% (range: -44% to 5%)
Didanosine 200 mg every 12 hours simultaneously administered with	1000 mg every 8 hours	12	No effect on ganciclovir PK parameters observed
ganciclovir	IV ganciclovir 5 mg/kg twice daily	11	No effect on ganciclovir PK parameters observed
	IV ganciclovir 5 mg/kg once daily	11	No effect on ganciclovir PK parameters observed
Trimethoprim 200 mg once daily	1000 mg every 8 hours	12	Ganciclovir renal clearance ↓ 16.3% Half-life ↑15%

Table 12 Results of Drug Interaction Studies With Ganciclovir: Effects of Ganciclovir on Pharmacokinetic Parameters of Coadministered Drug

Coadministered Drug	Ganciclovir Dosage	N	Coadministered Drug Pharmacokinetic (PK) Parameter
Zidovudine 100 mg every 4 hours	1000 mg every 8 hours	12	AUC ₀₋₄ \uparrow 19 ± 27% (range: -11% to 74%)
Mycophenolate Mofetil (MMF) 1.5 g single dose	IV ganciclovir 5 mg/kg single dose	12	No PK interaction observed (patients with normal renal function)
Didanosine 200 mg every 12 hours when administered 2 hours prior to or concurrent with ganciclovir	1000 mg every 8 hours	12	AUC ₀₋₁₂ \uparrow 111 ± 114% (range: 10% to 493%)
Didanosine 200 mg every 12 hours	IV ganciclovir 5 mg/kg twice daily	11	AUC ₀₋₁₂ \uparrow 70 ± 40% (range: 3% to 121%) C _{max} \uparrow 49 ± 48% (range: -28% to 125%)
Didanosine 200 mg every 12 hours	IV ganciclovir 5 mg/kg once daily	11	AUC ₀₋₁₂ \uparrow 50 ± 26% (range: 22% to 110%) C _{max} \uparrow 36 ± 36% (range: -27% to 94%)
Trimethoprim 200 mg once daily	1000 mg every 8 hours	12	Increase (12%) in C _{min}

12.4 Virology

<u>Mechanism of action:</u> Valganciclovir is an L-valyl ester (prodrug) of ganciclovir that exists as a mixture of two diastereomers. After oral administration, both diastereomers are rapidly converted to ganciclovir by intestinal and hepatic esterases. Ganciclovir is a synthetic analogue of 2'-deoxyguanosine, which inhibits replication of human cytomegalovirus in cell culture and in vivo.

In CMV-infected cells ganciclovir is initially phosphorylated to ganciclovir monophosphate by the viral protein kinase, pUL97. Further phosphorylation occurs by cellular kinases to produce ganciclovir triphosphate, which is then slowly metabolized intracellularly (half-life 18 hours). As the phosphorylation is largely dependent on the viral kinase, phosphorylation of ganciclovir occurs preferentially in virus-infected cells. The virustatic activity of ganciclovir is due to inhibition of viral DNA synthesis by ganciclovir triphosphate.

Antiviral Activity: The quantitative relationship between the cell culture susceptibility of human herpes viruses to antivirals and clinical response to antiviral therapy has not been established, and virus sensitivity testing has not been standardized. Sensitivity test results, expressed as the concentration of drug required to inhibit the growth of virus in cell culture by 50% (EC₅₀), vary greatly depending upon a number of factors. Thus the EC₅₀ value of ganciclovir that inhibits human CMV replication in cell culture (laboratory and clinical isolates) has ranged from 0.08 to 22.94 μ M (0.02 to 5.75 μ g/mL). Ganciclovir inhibits mammalian cell proliferation (CIC₅₀) in cell culture at higher concentrations ranging from 40 to >1000 μ M (10.21 to >250 μ g/mL). Bone marrow-derived colony-forming cells are more sensitive (CIC₅₀ = 2.7 to 12 μ M (0.69 to 3.06 μ g/mL).

<u>Viral Resistance</u>: Viruses resistant to ganciclovir can arise after prolonged treatment with valganciclovir by selection of mutations in either the viral protein kinase gene (UL97) responsible for ganciclovir monophosphorylation and/or in the viral DNA polymerase gene (UL54). Virus with mutations in the UL97 gene is resistant to ganciclovir alone, whereas virus with mutations in the UL54 gene may show cross-resistance to other antivirals that target the same sites on viral DNA polymerase.

The current working definition of CMV resistance to ganciclovir in cell culture assays is EC_{50} value $\geq 6.0 \,\mu\text{M}$ ($\geq 1.5 \,\mu\text{g/mL}$). CMV resistance to ganciclovir has been observed in individuals (immunocompromized and neonates) receiving prolonged treatment with ganciclovir or valganciclovir. The possibility of viral resistance should be considered in patients who show poor clinical response or experience persistent viral excretion during therapy.

13 NONCLINICAL TOXICOLOGY

13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

No long-term carcinogenicity studies have been conducted with Valcyte. However, upon oral administration, valganciclovir is rapidly and extensively converted to ganciclovir. Therefore, like ganciclovir, valganciclovir is a potential carcinogen.

Ganciclovir was carcinogenic in the mouse at oral doses that produced exposures approximately $0.1\times$ and $1.4\times$, respectively, the mean drug exposure in humans following the recommended intravenous dose of 5 mg/kg, based on area under the plasma concentration curve (AUC) comparisons. At the higher dose there was a significant increase in the incidence of tumors of the preputial gland in males, forestomach (nonglandular mucosa) in males and females, and reproductive tissues (ovaries, uterus, mammary gland, clitoral gland and vagina) and liver in females. At the lower dose, a slightly increased incidence of tumors was noted in the preputial and harderian glands in males, forestomach in males and females, and liver in females. Ganciclovir should be considered a potential carcinogen in humans.

Valganciclovir increases mutations in mouse lymphoma cells. In the mouse micronucleus assay, valganciclovir was clastogenic. Valganciclovir was not mutagenic in the Ames Salmonella assay. Ganciclovir increased mutations in mouse lymphoma cells and DNA damage in human lymphocytes in vitro. In the mouse micronucleus assay, ganciclovir was clastogenic. Ganciclovir was not mutagenic in the Ames Salmonella assay.

Valganciclovir is converted to ganciclovir and therefore is expected to have similar reproductive toxicity effects as ganciclovir [see Warnings and Precautions (5.2)]. Ganciclovir caused decreased mating behavior, decreased fertility, and an increased incidence of embryolethality in female mice following intravenous doses that produced an exposure approximately 1.7× the mean drug exposure in humans following the dose of 5 mg/kg, based on AUC comparisons. Ganciclovir caused decreased fertility in male mice and hypospermatogenesis in mice and dogs following daily oral or intravenous administration. Systemic drug exposure (AUC) at the lowest dose showing toxicity in each species ranged from 0.03 to 0.1x the AUC of the recommended human intravenous dose. Valganciclovir caused similar effects on spermatogenesis in mice, rats, and dogs. It is considered likely that ganciclovir (and valganciclovir) could cause inhibition of human spermatogenesis.

13.3 Reproductive and Developmental Toxicology

Valganciclovir is converted to ganciclovir and therefore is expected to have reproductive toxicity effects similar to ganciclovir. Ganciclovir has been shown to be embryotoxic in rabbits and mice following intravenous administration, and teratogenic in rabbits. Fetal resorptions were present in at least 85% of rabbits and mice administered doses that produced 2× the human exposure based on AUC comparisons (all dose comparisons presented are based on the human AUC following administration of a single 5 mg/kg infusion of intravenous ganciclovir). Effects observed in rabbits included: fetal growth retardation, embryolethality, teratogenicity and/or maternal toxicity. Teratogenic changes included cleft palate, anophthalmia/microphthalmia, aplastic organs (kidney and pancreas), hydrocephaly and brachygnathia. In mice, effects observed were maternal/fetal toxicity and embryolethality. Daily intravenous doses administered to female mice prior to mating, during gestation, and during lactation caused hypoplasia of the testes and seminal vesicles in the month-old male offspring, as well as pathologic changes in the nonglandular region of the stomach [see Warnings and Precautions (5.3)]. The drug exposure in mice as estimated by the AUC was approximately 1.7× the human AUC. Data obtained using an ex vivo human placental model show that ganciclovir crosses the placenta and that simple diffusion is the most likely mechanism of transfer. The transfer was not saturable over a concentration range of 1 to 10 mg/mL and occurred by passive diffusion.

14 CLINICAL STUDIES

14.1 Adult Patients

Induction Therapy of CMV Retinitis: In one randomized open-label controlled study, 160 patients with AIDS and newly diagnosed CMV retinitis were randomized to receive treatment with either Valcyte tablets (900 mg twice daily for 21 days, then 900 mg once daily for 7 days) or with intravenous ganciclovir solution (5 mg/kg twice daily for 21 days, then 5 mg/kg once daily for 7 days). Study participants were: male (91%), White (53%), Hispanic (31%), and Black (11%). The median age was 39 years, the median baseline HIV-1 RNA was 4.9 log₁₀, and the median CD4 cell count was 23 cells/mm³. A determination of CMV retinitis progression by the masked review of retinal photographs taken at baseline and Week 4 was the primary outcome measurement of the 3-week induction therapy. **Table 13** provides the outcomes at 4 weeks.

Table 13 Week 4 Masked Review of Retinal Photographs in CMV Retinitis Study

	Intravenous Ganciclovir	Valcyte Tablets
Determination of CMV retinitis progression at Week	N=80	N=80
4		
Progressor	7	7
Non-progressor	63	64
Death	2	1
Discontinuations due to Adverse Events	1	2
Failed to return	1	1
CMV not confirmed at baseline or no interpretable	6	5
baseline photos		

Maintenance Therapy of CMV Retinitis: No comparative clinical data are available on the efficacy of Valcyte tablets for the maintenance therapy of CMV retinitis because all patients in the CMV retinitis study received open-label Valcyte tablets after Week 4. However, the AUC for ganciclovir is similar following administration of 900 mg Valcyte tablets once daily and 5 mg/kg intravenous ganciclovir once daily. Although the ganciclovir C_{max} is lower following Valcyte tablets administration compared to intravenous ganciclovir, it is higher than the C_{max} obtained following oral ganciclovir administration [see Figure 1 in Clinical Pharmacology (12.3)]. Therefore, use of Valcyte tablets as maintenance therapy is supported by a plasma concentration-time profile similar to that of two approved products for maintenance therapy of CMV retinitis.

Prevention of CMV Disease in Heart, Kidney, Kidney-Pancreas, and Liver Transplantation: A double blind, double-dummy active comparator study was conducted in 372 heart, liver, kidney, and kidney-pancreas transplant patients at high-risk for CMV disease (D+/R-). Patients were randomized (2 Valcyte: 1 oral ganciclovir) to receive either Valcyte tablets (900 mg once daily) or oral ganciclovir (1000 mg three times a day) starting within 10 days of transplantation until Day 100 post-transplant. The proportion of patients who developed CMV disease, including CMV syndrome and/or tissue-invasive disease during the first 6 months post-transplant was similar between the Valcyte tablets arm (12.1%, N=239) and the oral ganciclovir arm (15.2%, N=125). However, in liver transplant patients, the incidence of tissue-invasive CMV disease was significantly higher in the Valcyte group compared with the ganciclovir group. These results are summarized in **Table 14**.

Mortality at six months was 3.7% (9/244) in the Valcyte group and 1.6% (2/126) in the oral ganciclovir group.

Table 14 Percentage of Patients With CMV Disease and Tissue-Invasive CMV Disease by Organ Type: Endpoint Committee, 6 Month ITT Population

	CMV Disease*		Tissue-Invasive CMV Disease		CMV Syndrome	
Organ	VGCV	GCV	VGCV	GCV	VGCV	GCV
	(N=239)	(N=125)	(N=239)	(N=125)	(N=239)	(N=125)
Liver	19%	12%	14%	3%	5%	9%
(n=177)	(22 / 118)	(7 / 59)	(16 / 118)	(2 / 59)	(6 / 118)	(5 / 59)
Kidney	6%	23%	1%	5%	5%	18%
(n=120)	(5 / 81)	(9 / 39)	(1 / 81)	(2 / 39)	(4 / 81)	(7 / 39)
Heart	6%	10%	0%	5%	6%	5%
(n=56)	(2 / 35)	(2 / 21)	(0 / 35)	(1 / 21)	(2 / 35)	(1 / 21)
Kidney / Pancreas (n=11)	0%	17%	0%	17%	0%	0%
	(0 / 5)	(1 / 6)	(0 / 5)	(1 / 6)	(0 / 5)	(0 / 6)

GCV = oral ganciclovir; VGCV = valganciclovir

14.2 Pediatric Patients

Prevention of CMV in Pediatric Solid Organ Transplant Recipients: Sixty-three children, 4 months to 16 years of age, who had a solid organ transplant (kidney 33, liver 17, heart 12, and kidney/liver 1) and were at risk for developing CMV disease, were enrolled in an open-label, safety, and pharmacokinetic study of oral Valcyte (Valcyte for oral solution or tablets). Patients received Valcyte once daily as soon as possible after transplant until a maximum of 100 days post-transplant. The daily doses of Valcyte were calculated at each study visit based on body surface area and a modified creatinine clearance [see Dosage and Administration (2.3)].

^{*}Number of Patients with CMV Disease = Number of Patients with Tissue-Invasive CMV Disease + Number of Patients with CMV Syndrome.

The pharmacokinetics of ganciclovir were similar across organ transplant types and age ranges. The mean daily ganciclovir exposures in pediatric patients were comparable to those observed in adult solid organ transplant patients receiving Valcyte 900 mg once daily [see Clinical Pharmacology (12.3)]. No case of CMV disease was reported during the study. CMV viremia was reported in 7 (11%) patients during the study; however, none of these events fulfilled the definition of CMV syndrome. Based on the pharmacokinetic, safety, and efficacy data from this study and extrapolated efficacy data from the adult study, oral Valcyte is indicated for the prevention of CMV disease in kidney and heart transplant children 4 months to 16 years of age at risk for developing CMV disease. Valcyte is not approved in adults for CMV prophylaxis in liver transplant patients; therefore, Valcyte is not recommended for CMV prophylaxis in pediatric liver transplant patients because efficacy cannot be extrapolated from adults.

15 REFERENCES

- NIOSH Alert: Preventing occupational exposures to antineoplastic and other hazardous drugs in healthcare settings. 2004. U.S. Department of Health and Human Services, Public Health Service, Centers for Disease Control and Prevention, National Institute for Occupational Safety and Health, DHHS (NIOSH) Publication No. 2004-165.
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- American Society of Health-System Pharmacists. ASHP guidelines on handling hazardous drugs. Am J Health-Syst Pharm. 2006; 63:1172-1193.
- Polovich, M., White, J. M., & Kelleher, L.O. (eds.) 2005. Chemotherapy and biotherapy guidelines and recommendations for practice (2nd. ed.) Pittsburgh, PA: Oncology Nursing Society.

16 HOW SUPPLIED/STORAGE AND HANDLING

Valcyte Tablets

Supplied as 450 mg pink convex oval tablets with "VGC" on one side and "450" on the other side. Each tablet contains valganciclovir HCl equivalent to 450 mg valganciclovir. Valcyte is supplied in bottles of 60 tablets (NDC 0004-0038-22). Store at 25°C (77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) [See USP controlled room temperature].

Valcyte for Oral Solution

Supplied as a white to slightly yellow powder blend for constitution, forming a colorless to brownish yellow tutti-frutti flavored solution. Available in glass bottles containing approximately 100 mL of solution after constitution. Each bottle can deliver up to a total of 88 mL of solution. Each bottle is supplied with a bottle adapter and 2 oral dispensers (NDC 0004-0039-09).

Prior to dispensing to the patient, Valcyte for oral solution must be prepared by the pharmacist [see Dosage and Administration (2.4)]. Store dry powder at 25°C (77°F); excursions permitted to 15°C to 30°C (59°F to 86°F) [See USP controlled room temperature].

Store constituted solution under refrigeration at 2° C to 8° C (36° F to 46° F) for no longer than 49 days. Do not freeze.

17 PATIENT COUNSELING INFORMATION

See FDA-Approved Patient Labeling

Valcyte tablets cannot be substituted for ganciclovir capsules on a one-to-one basis. Inform patients switching from ganciclovir capsules of the risk of overdosage if they take more than the prescribed number of Valcyte tablets [see Dosage and Administration (2.1), Overdosage (10)].

Adult patients should use Valcyte tablets, not Valcyte for oral solution [see Dosage and Administration (2.5)].

Valcyte is changed to ganciclovir once it is absorbed into the body. Inform all patients that the major toxicities of ganciclovir include granulocytopenia (neutropenia), anemia, and thrombocytopenia and that dose modifications may be required, including discontinuation. The importance of close monitoring of blood counts while on therapy should be emphasized. Inform patients that ganciclovir has been associated with elevations in serum creatinine.

Instruct patients to take Valcyte with food to maximize bioavailability.

Advise patients that ganciclovir causes decreased sperm production in animals and may cause decreased fertility in humans. Advise women of childbearing potential that ganciclovir causes birth defects in animals and should not be used during pregnancy. Because of the potential for serious adverse events in nursing infants, instruct mothers not to breast-feed if they are receiving Valcyte. Advise women of childbearing potential to use effective contraception during and for at least 30 days following treatment with Valcyte. Similarly, advise men to practice barrier contraception during and for at least 90 days following treatment with Valcyte.

Although there is no information from human studies, advise patients that ganciclovir should be considered a potential carcinogen.

Convulsions, sedation, dizziness, ataxia and/or confusion have been reported with the use of Valcyte and/or ganciclovir. If they occur, tasks requiring alertness may be affected including the patient's ability to drive and operate machinery.

Inform patients that ganciclovir is not a cure for CMV retinitis, and they may continue to experience progression of retinitis during or following treatment. Advise patients to have ophthalmologic follow-up examinations at a minimum of every 4 to 6 weeks while being treated with Valcyte. Some patients will require more frequent follow-up.

FDA-Approved Patient Labeling

Read the Patient Information that comes with Valcyte (VAL-site) before you start using it and each time you get a refill. There may be new information. This information does not take the place of talking with your healthcare provider.

What is the most important information I should know about Valcyte?

- Valcyte can affect your blood cells and bone marrow causing serious and life-threatening problems. Valcyte can lower the amount of your white blood cells, red blood cells, and platelets. Your doctor may do regular blood tests to check your blood cells while you are taking Valcyte. Based on these tests, your doctor may change your dose or tell you to stop taking Valcyte.
- Valcyte may cause cancer. Valcyte causes cancer in animals. It is not known if Valcyte causes cancer in people.
- Valcyte may cause birth defects. Valcyte causes birth defects in animals. It is not known if Valcyte causes birth defects in people. Valcyte should not be used during pregnancy. Tell your doctor right away if you get pregnant while taking Valcyte. If you can get pregnant, you should use effective birth control during treatment with Valcyte and for at least 30 days after treatment. Men should use a condom during treatment with Valcyte, and for at least 90 days after treatment, if their partner can get pregnant. Talk to your doctor if you have questions about birth control. Valcyte may lower the amount of sperm in a man's body and cause fertility problems.
- Valcyte changes into the medicine ganciclovir once it is in your body. Ganciclovir is also the active ingredient in Cytovene EV and ganciclovir capsules. Do not take ganciclovir capsules or Cytovene-IV if you are taking Valcyte. The dose of medicine in Valcyte tablets and ganciclovir capsules is different. One tablet of Valcyte has more medicine than one capsule of ganciclovir. This means that one Valcyte tablet cannot be substituted for one ganciclovir capsule. You could overdose and become very sick if Valcyte is taken with ganciclovir capsules or Cytovene-IV. Talk to your doctor or pharmacist if you have questions about your medicine.

What is Valcyte?

Valcyte is an "antiviral" medicine used in:

Adults:

- to treat cytomegalovirus (CMV) retinitis in people who have acquired immunodeficiency syndrome (AIDS). When CMV virus infects the eyes, it is called CMV retinitis.
- to prevent cytomegalovirus (CMV) disease in people who have received a **heart**, **kidney**, **or kidney-pancreas** transplant and who have a chance for getting CMV disease.

Children (4 months to 16 years of age):

• to prevent cytomegalovirus (CMV) disease in children who have received a **heart or kidney** transplant and who have a chance for getting CMV disease.

Valcyte may

- slow the growth of CMV virus in your body. CMV is an infection caused by a herpes virus called cytomegalovirus. If CMV retinitis isn't treated, it can cause blindness. Valcyte may protect your eyesight from damage due to CMV disease. CMV can also infect other parts of the body.
- prevent CMV disease for up to 6 months after **heart**, **kidney**, **or kidney-pancreas** transplant. Valcyte may prevent CMV virus from spreading into healthy cells.

Valcyte does not cure CMV retinitis. You may still get retinitis or worsening of retinitis during or after treatment with Valcyte. Therefore, it is important to stay under a doctor's care and have your eyes checked regularly.

The safety and efficacy of Valcyte have not been established in pediatric patients younger than 4 months of age.

The safety and efficacy of Valcyte have not been studied in adults older than age 65.

Who should not take Valcyte?

- Do not take Valcyte tablets if you are receiving hemodialysis. The use of ganciclovir capsules rather than Valcyte tablets is recommended.
- Do not take Valcyte if you are allergic to any of its ingredients or if you have ever had a serious reaction to ganciclovir capsules or Cytovene-IV. See the end of this leaflet for a list of the ingredients in Valcyte.

In addition, Valcyte is not for use in prevention of CMV disease in adults or children who have received a liver transplant. More research is needed before Valcyte can be recommended for use in the prevention of CMV disease in organ transplant patients other than those indicated (such as lung transplant patients).

Before taking Valcyte, tell your doctor:

• if you are pregnant or plan to become pregnant. Valcyte may cause birth defects. (See "What is the most important information I should know about Valcyte?")

- if you are breast-feeding. It is not known if Valcyte passes into your milk and if it may harm your baby. You should not breast-feed if you are HIV-positive because of the chance of passing the HIV virus to your baby through your milk.
- if you have kidney problems. Your doctor may give you a lower dose of Valcyte, or check you more often if you are taking Valcyte.
- if you have blood cell problems
- if you are having radiation treatment
- about all the medicines you take, including prescription and non-prescription medicines, vitamins and herbal supplements. Do not take ganciclovir capsules or Cytovene-IV if you are taking Valcyte. Valcyte and other medicines may affect each other. These interactions may cause serious problems. The following medicines may need dose changes if you are also taking Valcyte:
- Videx[®] (didanosine, ddI)
- Retrovir[®] (zidovudine, ZDV, AZT)
- Probeneció

Tell your doctor if you take medicines such as chemotherapy medicines that can lower your bone marrow function.

How should I take Valcyte?

- Valcyte is taken as a tablet for adults and either an oral solution or tablet for children. Take Valcyte exactly as your doctor prescribes it.
- If your doctor prescribes you Valcyte for oral solution, your pharmacist will give you dosing dispensers to measure your dose of Valcyte for oral solution. In order to be sure you receive the recommended dose, it is important to use the dispenser provided to you. Review the instructions below on how to take Valcyte for oral solution and how to use the dispenser. Ask your pharmacist if you have any questions. If you lose or damage your dispensers and cannot use them, contact your healthcare professional or pharmacist.
- Your dose of Valcyte will depend on your medical condition. If you have kidney problems or you are over age 65, your doctor may give you a lower dose of Valcyte.
- the usual dose for adults to get active CMV retinitis under control (induction therapy) is two 450 mg tablets twice a day for 21 days.
- the usual dose for adults to help keep CMV retinitis under control (maintenance therapy) is two 450 mg tablets once a day.
- the usual dose to prevent CMV in adults who have had a **heart, kidney, or kidney-pancreas** transplant is two 450 mg tablets once a day starting within 10 days of transplant and continuing until 100 days after the transplant.
- Take Valcyte with food.
- Do not break or crush Valcyte tablets.
- If you miss a dose of Valcyte, take the missed dose as soon as you remember. Then, take the next dose at the usual scheduled time. However, if it is almost time for your next dose, **do not take the missed dose.**
- Do not let your Valcyte run out. The amount of virus in your blood may increase if your medicine is stopped, even for a short time.
- If you take too much Valcyte, call your local poison control center or emergency room right away. You may need treatment in a hospital.
- Do not substitute Valcyte tablets for ganciclovir capsules. Adult patients should use Valcyte tablets, not Valcyte for oral solution. Talk to your doctor, nurse or pharmacist if you have questions about your medicine.

What should I avoid while taking Valcyte?

- Do not get pregnant. Valcyte causes birth defects in animals. It is not known if Valcyte causes birth defects in people. Valcyte should not be used during pregnancy. Tell your doctor right away if you get pregnant while taking Valcyte. If you can get pregnant, you should use effective birth control during treatment with Valcyte and for at least 30 days after treatment. Men should use a condom during treatment with Valcyte, and for at least 90 days after treatment, if their partner can get pregnant. Talk to your doctor if you have questions about birth control. Valcyte may lower the amount of sperm in a man's body and cause fertility problems.
- **Do not breast-feed.** Valcyte may harm your baby. You should not breast-feed if you are HIV-positive because of the chance of passing the HIV virus to your baby through your milk.
- Do not drive a car or operate other dangerous machinery until you know how Valcyte affects you. Valcyte can cause seizures, sleepiness, dizziness, unsteady movements, and confusion.

• **Do not break or crush Valcyte tablets.** Avoid contact with broken Valcyte tablets on your skin, mucous membranes or eyes. If contact occurs, wash your skin well with soap and water or rinse your eyes well with plain water.

What are the possible side effects of Valcyte?

See "What is the most important information I should know about Valcyte?" for details on the most serious side effects. Valcyte can also cause the following serious side effects:

- **kidney problems.** Valcyte may affect your kidney function. Your doctor may do regular blood tests called serum creatinine levels to check your kidney function while you are taking Valcyte.
- brain and nerve problems. Valcyte may cause seizures, sleepiness, dizziness, unsteady movements, and confusion.

Common side effects of Valcyte include diarrhea, nausea, vomiting, stomach pain, fever, headache, shaky movements (tremors), graft rejection, swelling of the legs, constipation, back pain, trouble sleeping, and high blood pressure.

Common changes in blood tests for people taking Valcyte include low white blood cells (neutropenia or leukopenia), low red blood cells (anemia), increased blood creatinine levels, increased calcium in the blood, and abnormal liver function.

Talk to your doctor about side effects that bother you or that won't go away.

These are not all the side effects of Valcyte. For more information, ask your doctor or pharmacist.

How do I store Valcyte?

- Valcyte tablets should be stored at room temperature, 59° to 86°F (15° to 30°C).
- Valcyte for oral solution should be stored under refrigeration at 36° to 46°F (2° to 8°C), for no longer than 49 days. Do not freeze.
- Keep Valcyte and all medicines out of the reach of children. Do not keep medicine that is out of date or that you no longer need. Dispose of unused medicines through community take-back disposal programs when available or place Valcyte in an unrecognizable, closed container in the household trash.

General information about Valcyte

Medicines are sometimes prescribed for conditions that are not mentioned in patient information leaflets. Do not use Valcyte for a condition for which it was not prescribed. Do not give Valcyte to other people, even if they have the same symptoms you have. It may harm them.

This leaflet summarizes the most important information about Valcyte. If you would like more information, talk with your doctor. You can ask your doctor or pharmacist for information about Valcyte that is written for health professionals. Information about Valcyte is also available at 1-800-526-6367 (toll-free).

What are the ingredients in Valcyte?

Active Ingredient: valganciclovir HCl

Inactive Ingredients for Tablets: microcrystalline cellulose, povidone K-30, crospovidone, and stearic acid. The film-coating applied to the tablets contains Opadry Pink[®].

Inactive Ingredients for Oral Solution: sodium benzoate, fumaric acid, povidone K-30, sodium saccharin, mannitol and tutti-frutti flavoring.

Dosing Instructions for Patients

How do I take Valcyte for oral solution?

Valcyte for oral solution will be prepared by the pharmacist prior to dispensing the medication to you.

Please follow instructions carefully to ensure proper dosing of the oral solution.

Figure 1:



- Shake closed bottle well for about 5 seconds before each use.
- Remove the child-resistant bottle cap.
- Before inserting the tip of the oral dispenser into bottle adapter, push the plunger completely down toward the tip of the oral dispenser. Insert tip firmly into opening of the bottle adapter.
- Turn the entire unit (bottle and oral dispenser) upside down (see Figure 1).
- Pull the plunger out slowly until the desired amount of medication is withdrawn into the oral dispenser (see Figure 2). The 18 mL (900 mg) dose is obtained by filling the dispenser twice, once to the 8 mL (400 mg) graduation, and a second fill to the 10 mL (500 mg) graduation.

Figure 2:



- Turn the entire unit right side up and remove the oral dispenser slowly from the bottle.
- Dispense directly into mouth and swallow. Do not mix with any liquid prior to dispensing.
- Close bottle with child-resistant bottle cap after each use.
- Immediately after receiving your medication, disassemble oral dispenser, rinse under running tap water and air dry prior to next use.

Care should be taken to avoid contact of the skin with the solution. If such contact occurs, wash thoroughly with soap and water. Do not use the solution after the expiration date.

Contact your doctor or pharmacist if your oral dispenser is lost or damaged, and they will advise you on how to continue to take your medication.

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Retrovir is a registered trademark of GlaxoSmithKline.

Distributed by:

Roche Laboratories Inc.

340 Kingsland Street

Nutley, New Jersey 07110-1199

VECO_1079070_PI_042008_N(1)

Revised: August 2009

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Representative sample of labeling (see the HOW SUPPLIED section for complete listing):

PRINCIPAL DISPLAY PANEL - 450 mg - 60 Tablet Bottle

NDC 0004-0038-22

Valcyte[®]

(valganciclovir HCl tablets)

equivalent to 450 mg valganciclovir

450 mg

60 tablets

Rx only



PRINCIPAL DISPLAY PANEL - 50 mg / 1 mL Bottle

NDC 0004-0039-09

Valcyte[®]

(valganciclovir

hydrochloride)

for oral solution

50 mg/1 mL

Each mL of constituted

oral solution contains

50 mg valganciclovir

free base.

Rx only

100 mL (3.4 fl oz)



PRINCIPAL DISPLAY PANEL - 50 mg / 1 mL Bottle Carton

NDC 0004-0039-09

$\mathbf{Valcyte}^{\mathbb{R}}$

(valganciclovir hydrochloride) for oral solution

50 mg/1 mL

Each mL of constituted oral solution contains 50 mg valganciclovir free base.

Rx only

100 mL (3.4 fl oz)



Revised: 08/2009 Distributed by: Hoffmann-La Roche Inc